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(54) A method of preparing polymer granules

Verfahren zur Herstellung von Polymergranulaten

Procédé pour la préparation de granules de polymère

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Description**Field of the Invention**

5 [0001] This invention relates to a method for preparing granules comprising a water soluble or water dispersable polymer, in which the polymer is introduced into an essentially horizontal drum containing a multiplicity of rotating arms proximating the internal surface of the drum, in which drum the polymer is present in a liquid and formed into granules in a single drying and granulation stage and to a device for the preparation of the said granules.

Background of the Invention

[0002] A two-step method of the above-mentioned type is known from JP-A-03 143 604. According to JP-A-03 143 604, a polymer is introduced into the drum in powdered form, whereas water is introduced into said drum separately. The polymer is allowed to take up water within the drum to a content of 15 - 40 % by weight, leading to the formation of a gel. Said gel is cleaved by the rotating arms. In a second step, the thus formed polymer granules are dried.

15 [0003] Granulated polymers are widely used in formulations where large quantities of water are undesirable, e.g. fabric washing powder, dishwashing powder, dishwashing tablets, water softening powder and -tablets, wettable powders containing agrochemicals. Polymers may be included to function as binders, dispersants, calcium sequestrants, etc.

[0004] Water soluble or water dispersable polymers in granular form are also used for a number of other applications like e.g. water treatment compositions, pigment dispersants, mineral dispersants, cement additives, oil field additives, textile additive compositions, adhesives, cosmetics, paper additives etc. In the following of the specification however, reference will be made to detergents, although the granules prepared are not limited to said use.

[0005] Detergent compositions are inter alia used for mechanical dishwashing and fabric washing. Detergent compositions intended for mechanical dishwashing (also known as machine dishwashing or automatic dishwashing) as well as for fabric washing usually require the presence of a component capable of binding Ca-ions (and Mg-ions) present in the washing liquor, said component usually being referred to as a builder.

[0006] Ca-ions (and Mg-ions) can react with alkali metalsilicates, carbonates and soap to give a precipitate which is deposited onto the washed product, like dishes, fabrics etc., being undesirable.

[0007] Conventional builders comprise phosphate (e.g. sodium tripolyphosphate, generally referred to as STP or STPP) and well known organic acids like citric acid, succinic acid etc.

[0008] During the past all kinds of polymers have been increasingly used as co-builders enhancing the Ca and/or Mg binding performance.

[0009] In more recent mechanical dishwashing powders phosphate is being replaced for environmental reasons by the weaker organic acids or acid salts, preferably the sodium acid salt e.g. of citric acid. In modern formulations polymers are generally used to enhance the Ca and Mg binding properties of the builders. These polymers also prevent redeposition of soil/dirt on the product washed and are generally applied at levels of about 1-10 wt.% in mechanical dishwashing powders.

[0010] Mechanical dishwashing detergent products are nearly exclusively produced by dry mixing of the various components in granular form. As a result of this development there is a growing demand for the abovementioned polymers in granular form with suitable properties, for incorporation in the detergent composition.

[0011] Co-granules have been developed, consisting of two or more components required in mechanical dishwashing. Generally these co-granules are based on silicate, sodium carbonate or sodium bicarbonate in conjunction with an organic component comprising one or more of the polymers.

[0012] These granules however frequently lack one or more of homogeneity, suitable flow properties, strength, solubility, bulk density and morphology to meet the requirements of the detergent formulator. Furthermore the freedom of the formulator is restricted in selecting the optimum ratios in the detergent product.

[0013] These co-granules known in the art are generally produced in a two-stage process, starting with a solution containing the components, which is dried in a drying stage e.g. in a spray-drying tower, a ring dryer, a drum-type dryer etc. giving a fine powder which can be compacted in a compactor as available on the market. With respect to compaction reference is made to US-A-3 875 282 and US-A-3 931 036. The compacted product leaving the compactor as a thin sheet is ground, milled and classified to the desired particle size by suitable sieves. Oversized and fine particles can be recycled through the compactor.

[0014] A disadvantage of such a compaction process is that due to the low yield of the desired particle size (approximately 200-1250 µm for detergent powders) excessive recycling is generally necessary (frequently more than 50%). Further the process is a two-stage process, which is complicated and not economic.

[0015] Fabric washing compositions especially when based on a zeolite as the main builder, in particular zeolite 4A, also require the presence of polymers as discussed above. Due to the fact that these products are also more and more produced by dry mixing processes it is essential that ingredients for such fabric washing compositions are provided as

free flowing, non dusting granules of sufficient bulk density and solubility.

[0016] As a consequence also for fabric washing detergent powders there is an increasing demand for said polymers in granular form.

[0017] Polymers are usually used in fabric washing powders at levels of about 0.1-8 wt.%.

5 [0018] As the granules are generally transported pneumatically in the detergent composition producing factories, the particles should be sufficiently durable to minimize attrition, which causes fines and gives rise to dust problems. Further the polymer should preferably be sufficiently soluble or at least dispersable in water. It has been observed, that cogranules produced by compacting fine powders containing these polymers have a tendency to attract moisture leading to sticky products and to agglomeration.

10 [0019] The main object of the present invention is to provide a method for preparing granules of said polymers in a single stage, said granules being sufficiently soluble, having the desired bulk density and show a minimum attrition.

[0020] The invention therefore provides a method for preparing granules comprising a water soluble or water dispersable polymer, in which the polymer is introduced into an essentially horizontal drum containing a multiplicity of rotating arms proximating the internal surface of the drum, in which drum the polymer is present with a liquid and formed into 15 granules in a single drying and granulation stage, characterized in that the polymer is introduced into the drum as a solution or dispersion, that the drum has an elevated temperature and that a gas is introduced into the drum at an elevated temperature.

[0021] According to the invention excellent polymer granules are prepared in a single stage.

20 [0022] It is noted that the granules prepared according to the invention may also comprise other components which are useful for the specific application of the granules, e.g. for detergent composition components, like silicate components, surfactants etc., provided that these materials do not impede the granulation method.

[0023] In this respect reference is made to EP-A-0 385 956 and US-A-4 252 968.

EP-A-0 385 956 relates to a process for spheronization and a device for implementing said process. One or more powders are placed in a leaktight vessel having a substantially vertical axis comprising a bottom blade rotating about an 25 axis parallel to the axis of said vessel. Further a single or multiple spheronizing rotating tool is present with a substantially discoidal form. The powders are mixed and a binder solution is added, wherein the composition of powders and binder solution is spheronized by using the spheronizing tool. In a final stage the spheronized material can be dried by fluidisation. Drying of a polymer solution or dispersion in a substantially horizontal drum containing a multiplicity of rotating arms, proximate its internal surface is not disclosed.

30 [0024] US-A-4 252 968 discloses a continuous process for preparing granular polycarbonate comprising feeding polycarbonate powder to a tubular substantially vertical drier equipped with a rotating mechanical stirrer, in which the powder is fluidized by means of a co-current stream of inert heated gas or vapour. In this process again fluidisation is used instead of a drying operation in a substantially horizontal drum with rotating arms. Further in US-A-4 252 968 the rotational speed is relatively low.

35 [0025] Further reference is made to EP-A-0 526 978, wherein a method is disclosed for the preparation of alkali metal silicate granules in a single drying and granulation stage using a drum with a multiplicity of rotating arms.

[0026] Preferably the polymer is selected from the following polymers or a mixture thereof:

- polymers based on acrylic- or methacrylic acid, copolymers thereof, or mixtures thereof;
- polymers based on vinylacetate, including homopolymers, and copolymers with other monomers; and
- biopolymers.

40 [0027] The acrylic or methacrylic acid based polymers can be either homopolymers of these acids or mixtures thereof, copolymers with other monomers containing mono unsaturated carbon-carbon bonds (-enes) such as maleic acid, itaconic acid and vinylacetate maleic acid or of mixtures thereof. Preferred polymers are the homopolymers of acrylic- and methacrylic acid and the copolymers of these acids with maleic acid and/or mixtures thereof. These polymers can be supplied either in the acidic form or in a neutralized (or partially neutralized) form after neutralisation with an alkali, like an alkali metal hydroxide solution.

45 [0028] The (meth)acrylic/maleic ratio of the copolymer is not specifically limited. However a ratio comprised between 0.5:1 mole ratio of (meth)acrylic acid/maleic anhydride and 3.5:1 mole ratio of acrylic acid/maleic anhydride is preferred.

[0029] The vinylacetate based polymer include homopolymers of vinylacetate as well as copolymers with other monomers, containing mono-unsaturated carbon-carbon bonds, such as maleic acid and itaconic acid.

[0030] The biopolymers are preferably starch and starch-derivatives.

50 [0031] Concerning the acrylic- or methacrylic acid based polymers, it is noted that the polymers can vary in molecular weight and in composition depending on the ratio of the monomers of (meth)acrylic acid and maleic acid used during polymerisation and the degree of polymerisation. The polymers are soluble in water and can be neutralized partially or completely by addition of alkali for example an alkali metal hydroxide, preferably sodium hydroxide.

[0032] These polymers in acidic or neutral form are on the market and are offered by National Starch and Chemical

Company, Bridgewater, New Jersey (US) and National Starch and Chemical Speciality Polymers Limited, Braunston, Daventry, Northants (GB). Examples of such products as supplied by National Starch are:

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TRADE MARK:	PRODUCT
Alcosperse 475	a copolymer in acidic form
Alcosperse 175	a copolymer in neutralized form
Alcosperse 659	a homopolymer in acidic form
Alcosperse 602N	a homopolymer in neutralized form

- 15 [0033] In the method according to the invention a solution or dispersion of the polymer in a liquid is used. It is meant also to comprise emulsions and the like. Said liquid is not specifically limited, but is preferably a polar solvent and more preferably an aqueous solution or dispersion of the polymer is used.
- [0034] The solution or dispersion used advantageously comprises 20-60 % by weight, preferably 35 to 60 % by weight of the polymer and more preferably 40 to 50 % by weight thereof.
- 20 [0035] The gas introduced into the drum advantageously is an inert gas and preferably air is used. The gas serves as a drying medium for drying the polymer solution or dispersion and discharging the evaporated solvent.
- [0036] Preferably the polymer solution or dispersion is introduced into the upper region of the drum and the gas is introduced into the lower region of the drum. In this manner the single stage drying according to the invention is optimized.
- 25 [0037] It is essential, that the internal surface wall of the drum is heated, and preferably to a temperature between 80 °C and the decomposition temperature of the polymer, more preferably between 120 and 200 °C, and most preferably between 150 and 170 °C.
- [0038] The gas introduced is also heated, preferably to a temperature between 80 °C and 350 °C, more preferably between 120 and 250 °C, and most preferably between 150 and 200 °C.
- 30 [0039] It will be clear for an ordinary expert, that the specific temperatures of the wall and the gas are selected depending on the specific circumstances i.e. the solids content of the solution or dispersion, the temperature thereof, the dimensions of the drum, the rotational speed of the rotating arms etc.
- [0040] Said rotational speed of the arms is preferably controlled such that the tip speed of the arms is comprised between 15 and 28 m/s, preferably comprised between 20 and 24 m/s. The tip speed is the speed of the tips of the arms, which can be designed in the form of paddles, blades or ploughs or the like, relative to the inner surface of the drum.
- 35 [0041] The temperature of the polymer solution is of course not critical and can be selected from between ambient temperature to the boiling temperature thereof and for practical reasons a temperature between 50 and 90 °C is preferred.
- 40 [0042] With the method according to the invention granules with a wide range of particle sizes can be prepared. In particular the average particle size of the granules prepared is comprised between 0.1 and 2 mm, and preferably between 0.2 and 1.2 mm.
- [0043] In a specific embodiment of the method according to the invention the granules having an average particle size of below 0.2 mm and above 2 mm are recycled. Advantageously said granules having an average particle size of below 45 0.2 mm are recycled by introduction thereof into about the centre portion of the drum.
- [0044] By recycling the particles being too large and too small, a very efficient and economic method is obtained. The selection of the centre portion of the drum provides the additional advantage, that the particles do not have to be redissolved, before entering the drum, but can be introduced as dry particles, mixing with the material inside the drum which is formed into granules. The exact location of said introduction into the drum is not very critical, but the centre portion, i.e. seen in the transport direction of the material, is preferred.
- 50 [0045] The recycled particles can also be redissolved in the solution fed to the drum, or fed near or with the gas stream.
- [0046] With the method according to the present invention granules can be obtained comprising a water soluble or water dispersable polymer, suitable for use in detergents. Said granules preferably have a moisture content of below 55 15% by weight, and preferably between 5 and 10 % by weight. The solubility of the granules is preferably such that the granules dissolve as fast as possible, e.g. within 3 min in water of 20 °C, and preferably within 2 min. The ball mill friability of the granules is preferably below 10%.
- [0047] With the method according to the invention, the bulk density of the granules can be adjusted and said bulk

density is preferably comprised between 300 and 800 g/l.

[0048] The use of the granules according to the invention is not limited to detergent compositions, but these granules are also preferably used for a water treatment composition, a pigments dispersant, a mineral dispersant, a cement additive, an oil field additive, a textile additive composition, an adhesive, a paper additive, an agrochemical, or cosmetic formulation.

[0049] Finally a device is provided suitable for performing the method according to the invention at least comprising a tubular drum with heating means for heating the inner surface of the drum, a rotatable shaft with a multiplicity of arms proximate said inner surface, feed means for a solution or dispersion to be dried, feed means for drying-gas and discharge means for dried material and gas, characterized in that feed means are present near about the centre of the drum for introducing recycled dried material.

[0050] In the following the test methods are discussed as used to determine the properties of the granules obtained in the examples.

Test methods

[0051] **Bulk density:** A cylinder (height/diameter ratio of approximately 2, is filled with powder to a measured volume of 1 litre and the sample is weighed.

[0052] **Ball mill friability (BMF):** This measures the breakdown of granules under conditions representing high shear mixing. The sample of granulate is sieved to remove oversize ($>1200 \mu\text{m}$) and undersize ($<200\mu\text{m}$) and then split into two parts. One part is used to measure the size distribution by sieving. The other part is put in the ball mill.

[0053] The ball mill is a 10×10 cm cylinder, containing 50 porcelain balls of 1 cm diameter and operated at 90 rpm, while set at a declination of 16 degrees. After 5 minutes milling the sample is removed and sieved to determine the size distribution. Ball mill friability is expressed as the percentage increase in fines $< 200 \mu\text{m}$.

[0054] **Particle size:** Measured using standard sieves (Retsch).

[0055] **Moisture content:** 2 grams of material are transferred to a clean aluminium dish and put into a forced draught oven, maintained at a temperature of 130°C . Sample is kept in the oven for 1 hour, after which it is cooled and reweighed as quickly as possible. The weight loss represents the moisture content.

[0056] In the following the present invention will be illustrated by means of the accompanied drawings, in which:

- 30 Fig. 1 shows a vertical diagrammatic section through a granulation apparatus according to the invention;
- Fig. 2 shows a graph of the moisture pick-up v.s. the time of granules according to the invention and granules obtained by compaction; and
- Fig. 3 shows a graph of the conductivity v.s. the time of granules according to the invention and granules obtained by compaction.

Specific description of the invention.

[0057] An example of the method of the invention will now be given to illustrate but not limit the invention.

[0058] The apparatus used is shown in fig. 1 and comprises a double walled tubular drum 1 mounted horizontally. Heated oil was passed through the double wall cavity 2 to heat the inner surface of the drum to a required temperature. The drum had a length of 2 metres and a diameter of 0.35 metres. Along the cylindrical axis was positioned a rotatable shaft 3 having about 100 arms 4 equally spaced along its length with 4 arms fixed at each contact point. These arms have paddle ends which extend to be almost in contact with the inner wall. The shaft was rotated at 1200 rotations per minute being equal to a tip speed of 22 m/s, i.e. being the speed of the tips of the paddle ends.

[0059] An acrylic acid/maleic acid copolymer solution, having an acrylic/maleic ratio of approximately 3:1 and a mol. weight of approximately 70,000, at a density of approximately 1.3 kg/l (45% d.s.) was sprayed into the drum at an inlet 5 just above the axis at one side of the drum at a rate of about 70 kg/hour at a temperature of 60°C . Air heated to 190°C by an heat exchanger entered the drum at a rate of $600 \text{ m}^3/\text{h}$ at an inlet 6 next to the axis and close to the point where the polymer solution was introduced at the same end of the drum (i.e. co-current). The granular polymer product and air left the drum at the other end of the drum through conduit 7. The drum wall temperature was maintained at 160°C .

[0060] The polymer used was a copolymer of acrylic acid and maleic acid, Alcosperse 475 from National Starch and Chemical Company, Bridgewater New Jersey (U.S.).

[0061] The polymer solution was rapidly broken into droplets by the force of the air movement and then impacted against the drum wall by the rotational energy of the paddles. The droplets were continuously impacted with the wall as they progressed through the drum under the force of the air.

[0062] The polymer product was drawn off at the lowest point of the drum and was found to be substantially spherical particles having a particle size distribution of

50% below 250 micron
 35% 200 to 1,000 micron
 15% above 1,000 micron

5 [0063] The product did not stick to the drum wall or the moving parts and was free flowing. It had a water content of 8%, a bulk density of 480 g/l and a ball mill friability of below 5% for particles above 200 micron.

[0064] It was sieved to provide the desired product with a particle range of 0.2 mm to 1.2 mm after separation from the air and steam by means of a cyclone. The particles outside this range can be recycled by addition to the polymer solution.

10 [0065] In a second example the polymer solution was run under the same conditions. However fines were reintroduced into the apparatus via a dosing screw via 9. The weight ratio of fines to feed stream was 1:1. Instead of reintroducing the fines via the dosing screw 9, these can also be reintroduced via 8 however it appeared to be more efficient to reintroduce via 9. Therefore the feed 8 has been dotted.

15 [0066] The polymer product was drawn off at the lowest point of the drum and was found to be substantially spherical particles having a particle size distribution of

53% below 250 micron
 35% 200 to 1,000 micron
 12% above 1,000 micron

20 [0067] The product did not stick to the drum wall or the moving parts and was free flowing. It had a water content of 8%, a bulk density of 480 g/l and a ball mill friability of below 5% for particles above 200 micron.

[0068] It was sieved to provide the desired product with a particle range of 0.2 mm to 1.2 mm after separation from the air and steam by means of a cyclone. The particles below 0.2 mm were recycled via 9.

25 [0069] The above two examples were also performed with the polymer Alcosperse 175 of National Starch and Chemical Company, providing similar results.

[0070] In the following the moisture pick-up and the solubility of the granules according to the invention is compared with granules prepared by compacting a spray-dried powder.

30 [0071] Fig. 2 shows the moisture pick-up of granules as prepared according to the invention and described in the example, being compared to granules also prepared from Alcosperse 475, in which the same solution is spray-dried using a gas inlet temperature of 260-275 °C and a gas outlet temperature of 98-110 °C. Thereafter the dry powder with a maximum water content of 10% was compacted using the method according to US-A-3 875 282. The granules were sieved to provide the product with a particle range of 0.2-1.2 mm. Approximately 5 g of both samples of granules was weighed into an aluminium dish and placed in a humidity cabinet set at 78% R.H. and 38 °C. The sample was

35 reweighed at set times and the weight increase was determined as a percentage of the initial weight. The results are shown in fig. 2. The moisture pick-up of the example according to the invention and the comparative example according to the prior art were both determined twice. It is clear that the moisture pick-up of the granules of the invention is much lower than with the compacted granules.

40 [0072] For measuring the solution time of the same two granular products as described above 1.5 g powdered polymer (0.2-1.2 mm) was weighed. 500 ml deionised water was introduced into a bottle and was stirred. A conductivity electrode was inserted and the stirring speed was adjusted until a vortex just formed. The weighed powder was added to the stirred water and at the same time a stopwatch was started. The conductivity for each sample was measured at 10 s intervals up to 1 min and at 30 s intervals thereafter up to 3 min, or until two consecutive readings were the same. The result is plotted in the graph according to fig. 3. A standard potassium chloride solution was used as a control. It is

45 clear that the solubility of the polymer according to the invention is superior to the solubility of the polymer granules as prepared by compaction.

Claims

- 50 1. Method for preparing granules comprising a water soluble or water dispersable polymer, in which the polymer is introduced into an essentially horizontal drum containing a multiplicity of rotating arms proximating the internal surface of the drum, in which drum the polymer is present with a liquid and formed into granules in a single drying and granulation stage, characterised in that the polymer is introduced into the drum as a solution or dispersion, that the drum has an elevated temperature and that a gas is introduced into the drum at an elevated temperature.
- 55 2. Method according to claim 1, characterised in that the polymer is selected from the following polymers or a mixture thereof:

- polymers based on acrylic- or methacrylic acid, copolymers thereof, or mixtures thereof;
- polymers based on vinylacetate, including homopolymers and copolymers with other monomers; and
- biopolymers.

- 5 3. Method according to one or more of claims 1-2, **characterised in that** an aqueous solution or dispersion of the polymer is used.
- 10 4. Method according to one or more of claims 1-3, **characterised in that** the solution or dispersion used comprises 20 to 60 % by weight of the polymer.
- 15 5. Method according to claim 4, **characterised in that** the solution or dispersion used comprises 35 to 60 % by weight of the polymer.
- 15 6. Method according to claim 5, **characterised in that** the solution or dispersion used comprises 40 to 50 % by weight of the polymer.
- 15 7. Method according to one or more of claims 2-6, **characterised in that** the polymer comprises an (meth)acrylic homo-polymer, a copolymer of (meth)acrylic acid and maleic acid, or a mixture thereof.
- 20 8. Method according to one or more of the preceding claims, **characterised in that** an inert gas is used.
- 20 9. Method according to claim 8, **characterised in that** the inert gas used is air.
- 25 10. Method according to one or more of the preceding claims, **characterised in that** the polymer solution or dispersion is introduced into the upper region of the drum and the gas is introduced into the lower region of the drum.
- 25 11. Method according to one or more of the preceding claims, **characterised in that** the temperature of the internal surface of the drum is comprised between 80 °C and the decomposition temperature of the polymer.
- 30 12. Method according to claim 11, **characterised in that** the temperature of the internal surface of the drum is comprised between 120 and 200 °C.
- 30 13. Method according to claim 12, **characterised in that** the temperature of the internal surface of the drum is comprised between 150 and 170 °C.
- 35 14. Method according to one or more of the preceding claims, **characterised in that** the gas inlet temperature is comprised between 80 °C and 350 °C.
- 40 15. Method according to claim 14, **characterised in that** the gas inlet temperature is comprised between 120 and 250 °C.
- 40 16. Method according to claim 15, **characterised in that** the gas inlet temperature is comprised between 150 and 200 °C.
- 45 17. Method according to one or more of the preceding claims, **characterised in that** the rotational speed of the arms is controlled such that the tip speed of the arms is comprised between 15 and 28 m/s.
- 45 18. Method according to claim 17, **characterised in that** the tip speed of the arms is comprised between 20 and 24 m/s.
- 50 19. Method according to one or more of the preceding claims, **characterised in that** the average particle size of the granules prepared is comprised between 0.1 and 2 mm.
- 55 20. Method according to claim 19, **characterised in that** the average particle size of the granules prepared is comprised between 0.2 and 1.2 mm.
- 55 21. Method according to one or more of the preceding claims, **characterised in that** the granules having an average particle size of below 0.2 mm and above 2 mm are recycled.

22. Method according to one or more of the preceding claims, characterised in that the granules having an average particle size of below 0.2 mm are recycled by introduction thereof into about the longitudinal centre portion of the drum.
- 5 23. Device suitable for performing the method according to one or more claims 1-22, at least comprising a tubular drum (1) with heating means for heating the inner surface of the drum (1), a rotatable shaft (3) with a multiplicity of arms (4) proximate said inner surface, feed means (5) for a solution or dispersion to be dried, feed means for drying-gas and discharge means for dried material and gas, characterised in that feed means (9) are present near about the longitudinal centre of the drum for introducing recycled dried material.

10 **Patentansprüche**

1. Verfahren zur Herstellung von Körnern, umfassend ein wasserlösliches oder wasserdispersierbares Polymer, bei dem das Polymer in eine im wesentlichen horizontale Trommel mit einer Vielzahl von rotierenden Armen eingegeben wird, die bis nahe an die innere Oberfläche der Trommel heranreichen, wobei das Polymer in der Trommel mit einer Flüssigkeit vorliegt und zu Körnern in einem einzigen Trocknungs- und Granulierungsschritt umgewandelt wird, dadurch gekennzeichnet, daß das Polymer als Lösung oder Dispersion in die Trommel eingegeben wird, daß die Trommel eine erhöhte Temperatur aufweist und daß ein Gas bei erhöhter Temperatur in die Trommel eingeführt wird.
2. Verfahren nach Anspruch 1, dadurch gekennzeichnet, daß das Polymer ausgewählt wird aus den folgenden Polymeren oder einer Mischung davon:
- Polymere auf Basis von Acryl- oder Methacrylsäure, Copolymere davon, oder Mischungen davon;
 - Polymere auf Basis von Vinylacetat, einschließlich Homopolymere und Copolymere mit anderen Monomeren und
 - Biopolymere.
3. Verfahren nach einem oder mehreren der Ansprüche 1 bis 2, dadurch gekennzeichnet, daß eine wäßrige Lösung oder Dispersion des Polymers eingesetzt wird.
4. Verfahren nach einem oder mehreren der Ansprüche 1 bis 3, dadurch gekennzeichnet, daß die verwendete Lösung oder Dispersion 20 bis 60 Gew.% des Polymers enthält.
5. Verfahren nach Anspruch 4, dadurch gekennzeichnet, daß die verwendete Lösung oder Dispersion 35 bis 60 Gew.% des Polymers enthält.
6. Verfahren nach Anspruch 5, dadurch gekennzeichnet, daß die verwendete Lösung oder Dispersion 40 bis 50 Gew.% des Polymers enthält.
7. Verfahren nach einem oder mehreren der Ansprüche 2 bis 6, dadurch gekennzeichnet, daß das Polymer ein (Meth)acrylhomonopolymer, ein Copolymer von (Meth)acrylsäure und Maleinsäure, oder einem Gemisch davon enthält.
8. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß ein Inertgas verwendet wird.
9. Verfahren nach Anspruch 8, dadurch gekennzeichnet, daß das verwendete Inertgas Luft ist.
10. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Polymerlösung oder Dispersion in den oberen Teil der Trommel und das Gas in den unteren Teil der Trommel eingegeben wird.
11. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Temperatur der inneren Oberfläche der Trommel zwischen 80 °C und der Zersetzungstemperatur des Polymers liegt.
12. Verfahren nach Anspruch 11, dadurch gekennzeichnet, daß die Temperatur der inneren Oberfläche der Trommel zwischen 120 und 200 °C liegt.

13. Verfahren nach Anspruch 12, dadurch gekennzeichnet, daß die Temperatur der inneren Oberfläche der Trommel zwischen 150 und 170 °C liegt.
- 5 14. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Gaseinlaßtemperatur zwischen 80 °C und 350 °C liegt.
- 15 15. Verfahren nach Anspruch 14, dadurch gekennzeichnet, daß die Gaseinlaßtemperatur zwischen 120 und 250 °C liegt.
- 10 16. Verfahren nach Anspruch 15, dadurch gekennzeichnet, daß die Gaseinlaßtemperatur zwischen 150 und 200 °C liegt.
- 15 17. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Rotationsgeschwindigkeit der Arme so eingestellt wird, daß die Geschwindigkeit der Spitzen der Arme zwischen 15 und 28 m/s liegt.
18. Verfahren nach Anspruch 17, dadurch gekennzeichnet, daß die Geschwindigkeit der Spitzen der Arme zwischen 20 und 24 m/s liegt.
- 20 19. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die durchschnittliche Teilchengröße der hergestellten Körner zwischen 0,1 und 2 mm liegt.
- 25 20. Verfahren nach Anspruch 19, dadurch gekennzeichnet, daß die durchschnittliche Teilchengröße der hergestellten Körner zwischen 0,2 und 1,2 mm liegt.
21. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Körner mit einer durchschnittlichen Teilchengröße unter 0,2 mm und über 2 mm recycelt werden.
- 30 22. Verfahren nach einem oder mehreren der vorangehenden Ansprüche, dadurch gekennzeichnet, daß die Körner mit einer durchschnittlichen Teilchengröße unter 0,2 mm recycelt werden, indem sie etwa in den mittleren, sich in Längsrichtung erstreckenden Teil der Trommel eingegeben werden.
- 35 23. Vorrichtung zur Durchführung des Verfahrens gemäß einem oder mehreren der Ansprüche 1 bis 22, umfassend zumindest eine rohrförmige Trommel (1) mit einer Heizvorrichtung zum Erhitzen der inneren Oberfläche der Trommel (1), einer rotierenden Welle (3) mit einer Vielzahl von Armen (4), die nahe an die genannte innere Oberfläche heranreichen, Einspeisevorrichtungen (5) für eine zu trocknende Lösung oder Dispersion, Einspeisevorrichtungen für Trocknungsgas und Entladevorrichtungen für getrocknetes Material und Gas, dadurch gekennzeichnet, daß Einspeisevorrichtungen (9) etwa in der Nähe des sich in Längsrichtung erstreckenden Zentrums der Trommel zur Eingabe von rezykliertem getrocknetem Material vorhanden sind.

40 **Revendications**

1. Méthode de préparation de granules comprenant un polymère soluble dans l'eau ou qui peut être dispersé dans l'eau, dans laquelle le polymère est introduit dans un tambour pratiquement horizontal contenant une pluralité de bras qui tournent à proximité de la surface interne du tambour, dans lequel le polymère est présent avec un liquide et prend la forme de granules dans une seule phase de séchage et de granulation, caractérisée en ce que le polymère est introduit dans le tambour en solution ou dispersion, en ce que le tambour a température élevée et en ce qu'un gaz est introduit dans le tambour à température élevée.
- 45 2. Méthode suivant la revendication 1, caractérisée en ce que le polymère est choisi parmi les polymères suivants ou dans un mélange de ceux-ci:
- 50 - polymères basés sur de l'acide acrylique ou méthacrylique, des copolymères de ceux-ci ou des mélanges de ceux-ci;
- 55 - polymères basés sur de l'acétate de vinyle, y compris des homopolymères et des copolymères avec d'autres monomères; et
- biopolymères.

3. Méthode suivant une des revendications 1 et 2, caractérisée en ce qu'on utilise une solution ou une dispersion aqueuse du polymère.
4. Méthode suivant une ou plusieurs des revendications 1 à 3, caractérisée en ce que la solution ou la dispersion utilisée comprend 20 à 60 % en poids du polymère.
5. Méthode suivant la revendication 4, caractérisée en ce que la solution ou la dispersion utilisée comprend 35 à 60 % en poids du polymère.
10. Méthode suivant la revendication 5, caractérisée en ce que la solution ou la dispersion utilisée comprend 40 à 60 % en poids du polymère.
15. Méthode suivant une ou plusieurs des revendications 2 à 6, caractérisée en ce que le polymère comprend un homo-polymère (méth)acrylique, un copolymère d'acide (méth)acrylique et d'acide maléique, ou un mélange de ceux-ci.
20. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce qu'on utilise un gaz inerte.
25. Méthode suivant la revendication 8, caractérisée en ce que le gaz inerte utilisé est de l'air.
30. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que la solution ou la dispersion de polymère est introduite dans la région supérieure du tambour et en ce que la gaz est introduit dans la région inférieure du tambour.
35. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que la température de la surface interne du tambour est comprise entre 80° C et la température de décomposition du polymère.
40. Méthode suivant la revendication 11, caractérisée en ce que la température de la surface interne du tambour est comprise entre 120 et 200° C.
45. Méthode suivant la revendication 12, caractérisée en ce que la température de la surface interne du tambour est comprise entre 150 et 170° C.
50. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que la température d'entrée du gaz est comprise entre 80 et 350°C.
55. Méthode suivant la revendication 14, caractérisée en ce que la température d'entrée du gaz est comprise entre 120 et 250° C.
60. Méthode suivant la revendication 15, caractérisée en ce que la température d'entrée du gaz est comprise entre 150 et 200° C.
65. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que la vitesse de rotation des bras est commandée de façon que la vitesse du bout des bras soit comprise entre 15 et 28 m/s.
70. Méthode suivant la revendication 17, caractérisée en ce que la vitesse du bout des bras est comprise entre 20 et 24 m/s.
75. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que la taille moyenne des particules des granules préparées est comprise entre 0,1 et 2 mm.
80. Méthode suivant la revendication 19, caractérisée en ce que la taille moyenne des particules de granules préparées est comprise entre 0,2 et 1,2 mm.
85. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que les granules qui ont une taille moyenne de particules inférieure à 0,2 mm et supérieure à 2 mm sont recyclées.
90. Méthode suivant une ou plusieurs des revendications précédentes, caractérisée en ce que les granules qui ont une

EP 0 734 763 B1

taille moyenne de particules inférieure à 0,2 mm sont recyclées par introduction de celles-ci dans à peu près la partie centrale longitudinale du tambour.

- 5 23. Dispositif qui convient pour mettre en oeuvre la méthode suivant une ou plusieurs des revendications 1 à 22, comprenant au moins un tambour tubulaire (1) avec des moyens de chauffage pour chauffer la surface interne du tambour (1), un axe rotatif (3) avec une multiplicité de bras (4) à proximité de ladite surface interne, des moyens d'alimentation (5) d'une solution ou d'une dispersion à sécher, des moyens d'alimentation en gaz séchant et des moyens d'évacuation du matériau séché et du gaz, caractérisé en ce que des moyens d'alimentation (9) sont prévus dans la région du centre longitudinal du tambour pour introduire le matériau recyclé séché.

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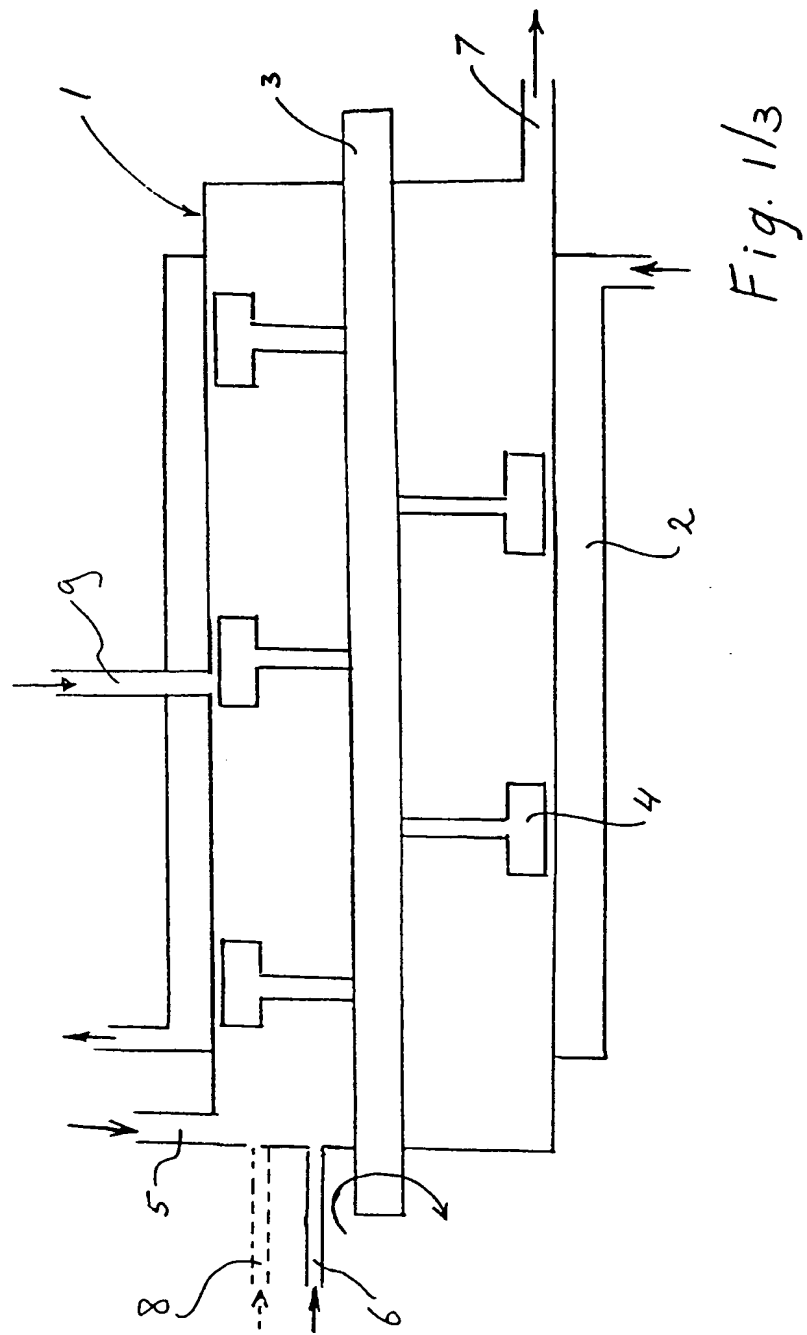


Fig. 1/3

% MOISTURE PICK-UP

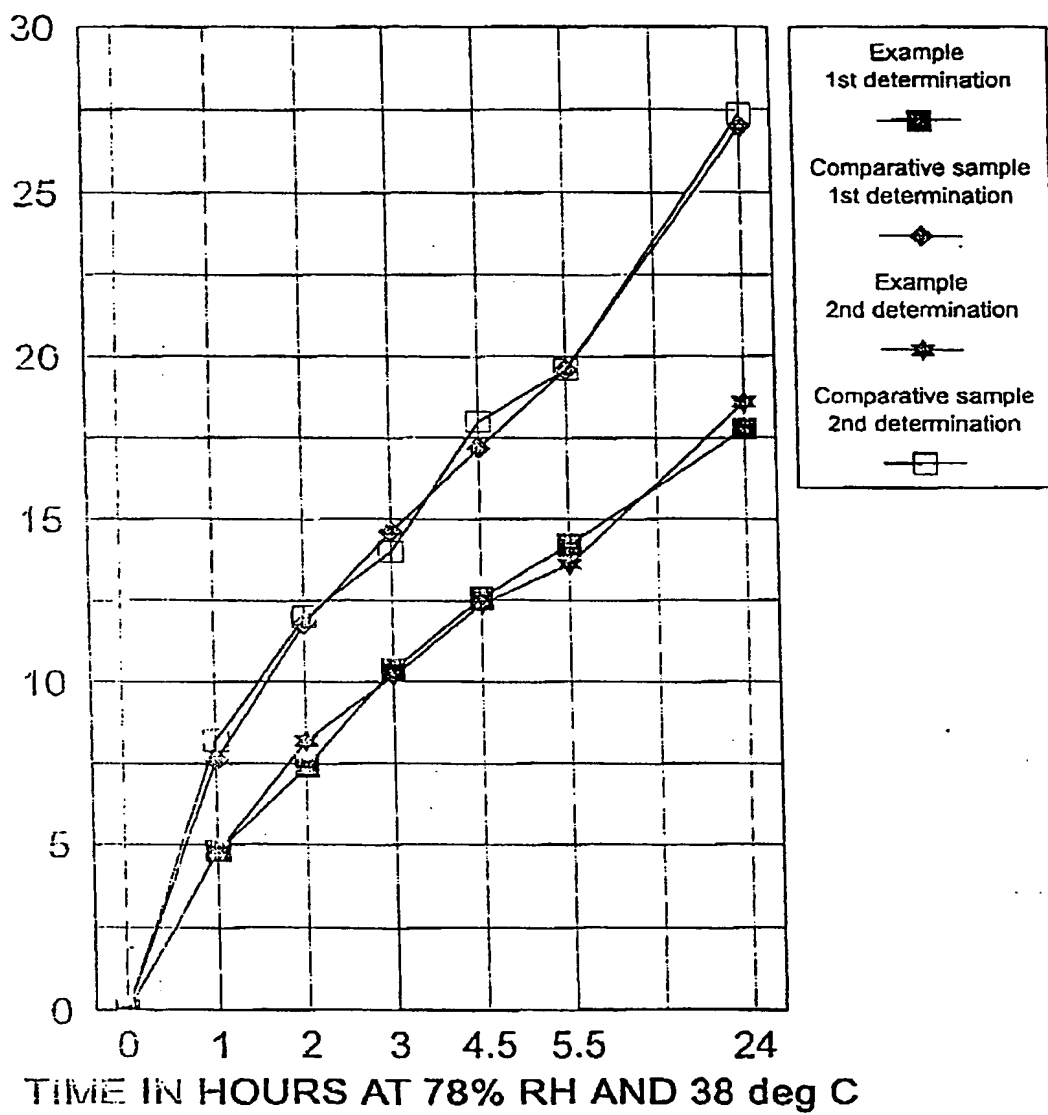


Fig. 2/3

CONDUCTIVITY (microS/cm)

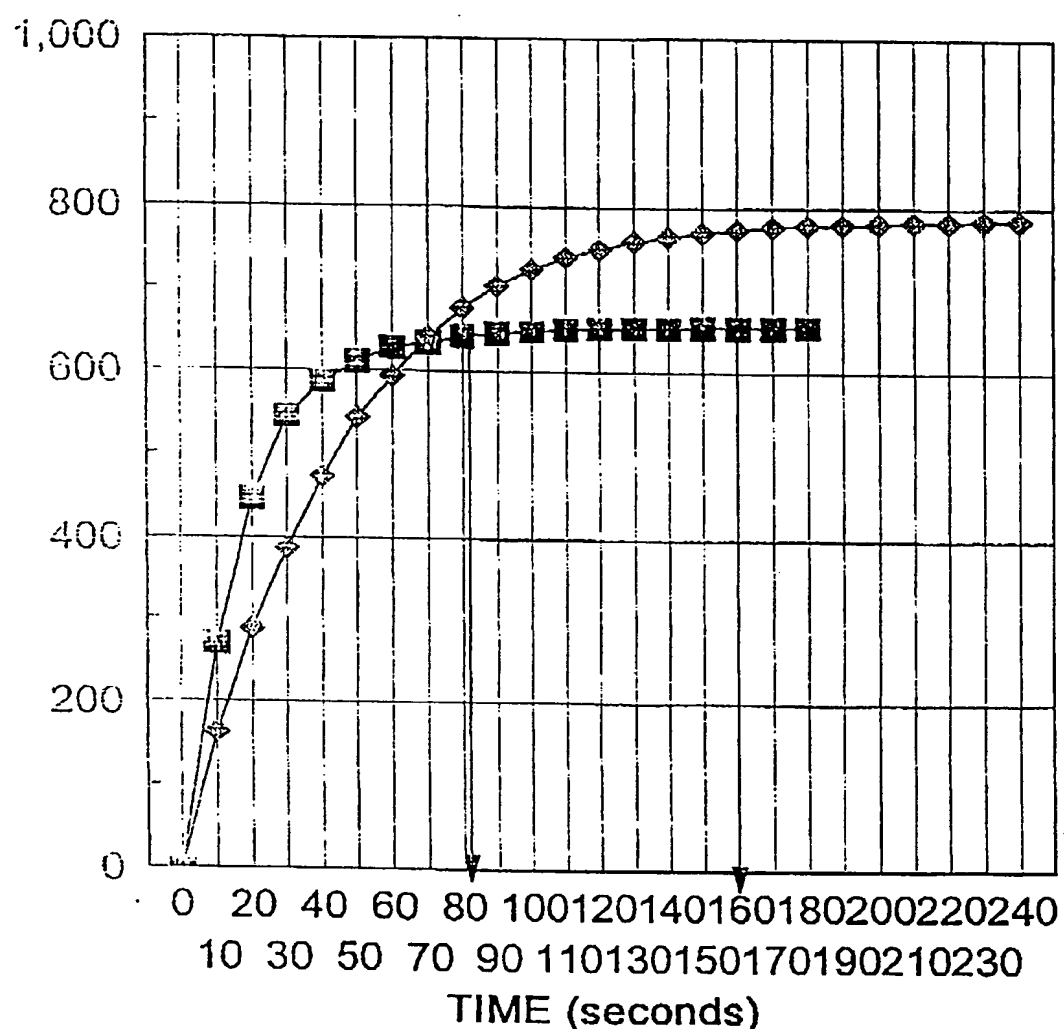


Fig.3/3

